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- (54) A CATALYTIC VAPOR GROWTH METHOD FOR PRODUCING CARBON FIBRILS

EIN KATALYTISCHES DAMPFZUCHTVERFAHREN ZUR HERSTELLUNG VON KOHLENSTOFFIBRILLEN

PROCEDE DE CRISTALLISATION EN PHASE VAPEUR PAR VOIE CATALYTIQUE POUR LA PRODUCTION DE FIBRILLES DE CARBONE

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The file contains technical information submitted after the application was filed and not included in this specification

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## Description

[0001] This application refers to fibrits. It more particularly refers to carbon/graphite fibrits and to an improved process for producing such. Carbon fibrits as used herein means graphitic fibrits having keys surface area, high young's modulus of elasticity and high tensile strength which are grown catalytically from available sources of carbon.

## Background of the Invention

[0002] It has been known for some time that one could make fibrilis by decomposing various carbon contributing molecules, such as light hydrocarbons, in contact with a suitable metal catalyst, such as for example ron alone or in combination with other metals. PCT Patent Publication WO 87/07599 to Tennent for example.

relates to the production of graphitic carbon fibritis.

(0003) More particularly the invention refers to an animproved process for producing such carbon/graphite fibritis. Fibritis are made according to this invention in a high temperature, catalytic process. The fibrit carb be made of a variety of materials, e.g. carbon, silicon nitride, silicon carbide, etc. Such fibritis have the atoms as in their composition relatively ordered at their outer surricess as they are made by this process. Thus, if carb the said that this process preferably directly produces a product having a relatively crystalline outer region to substantial portions of its length and may have inner or egions where its atoms are less ordered. It may, and often dose, even have a hollow region adality positioned along substantial portions of its length.

(0004) Fibrils prepared by a process according to this invention are characterized by small dismeters, e.g. 3.5 3 or 70 namenters and high LD up to about 100 and even more. Where the preferred structure described above is produced, it is exitably produced directly in the fixell forming process without further processing being required.

[0005] According to the present invention, Carbon fibrils can be produced in quite high yields in this embodiment, a suitable source of carbon may be a hydrocarbonacous material illustrated by methane, by hydrocarbonacous material illustrated by methane, propane, butane, benzene, cyclohoxane, 45 butene, isobutene, ethylene, propylene, acetylene, but-butene, isobutene, ethylene, propylene, acetylene, but-butene, isobutene, ethylene, promylene, acetylene, but-butene, isobutene, ethylene, chiralichyde, acetolae, hydrocarbonacous materials) other similar materials on-hydrocarbonacous materials) other similar materials, and mixtures of two (2) or more thereof Such feed is contacted with a suitable, carbon site of such side of the suitable, carbon fibrils to cause organistic carbon fibrils to orwor.

[0006] It is within the scope of this invention to provide a non-hydrocarbonaceous gas along with the carbon contributing reactant. Such gas might for example be hydrogen or carbon monoxide. Inert difuents are also

#### cuitable

[0007] The temperature of the process of this invention can vary widely depending upon the nature of the carbon source being used, however, the temperature is kept below the thermal decomposition temperature thereot. In the case of using a mixture of such carbon sources, the operating temperature should be maintained below the termal decomposition temperature of the most temperature should be maintained below the termal decomposition temperature of the most temperature in the range of 500 to 1500°C may be found to be generally usable, depending on the carbon source used, preferably between about 600 and opport.

19088] Subatmospheric, attmospheric ant/or superatmospheric pressures may be used as clicated by other other processing considerations. It has been found that it is desirable to provide the cathon source in the vapor state, and thus, the pressure should not be so high as to cause the carbon source to be in the liquid state under that forming temperature conditions. Further, it is desirable atthough not essential to provide a suitable gazeous diluent, such as hydrogen or inert gases, for example, nitrogen.

[0099] It is preferred that the system as a whole be non-oddizing wheretor preferably avoiding the presence of oxygen if practical. Small amounts of these materials can be tolerated. It should be understood that the existence of oxidizing conditions, at the elevated temperatures operative for this process, will cause oxidation of the carbon source and therefor reduce the amount of carbon from such source which is available for conversion into fixing as desired.

(0010) It may be desirable to provide suitable heat to this reaction system where and when needed. Temporature of different parts of the reactor zone may be suitably controlled to different temperatures and this is easily accomplished by using electrical resistance heating. However in larger scale industrial practices, electric resistance heating may sometimes be economically replaced by direct heating, such as for example by burning some of the carbon contributing feed to raise the temperature of the remainder of the feed, or by feeding the catalyst or the carbon contributing feed or the dilucent into the system at a sufficiently elevated temperature such that direct heat exchange of the component with each other will cause the fibril forming reaction to proceed as designed.

[0011] The nature of the catalyst seams to have a significant effect upon the yield of fibrilis produced according to this invention. It is known to use iron group metals such as iron, cobatt or nickel to catalyze the conversion of carbon contributing compounds to fibrilis, and such metals are within the scope of this invention. In addition, many other multivalant transition metals, including janthanides, appear to be operative. Particularly useful catalyte metals include inter aliae: iron, molydenum, cobatt, nickel, platinum, palladium, variandium, and chromium. The present process uses catalyst particles commium. The present process uses catalyst particles comprising as the multivalent metal, iron mixed with at least one other transition metal.

[0012] Particularly useful combinations include iron and molybdenum, iron and chromium, copper and inickel, iron and platinum, iron and tin, iron and nickel, siron and manganese, and iron and cerium.

[0013] The yield of fibrile produced according to the practice of this invention appears to be related to the physical state of the catalyst used to produce such. According to the invention, the multivalent transition metal fibril forming catalyst is present on a suitable substate as relatively discrete catalytic sites, each 3.5 to 70nm (35 to 700Å), preferably 6 to 30nm (60 to 300Å) in size during fibril formation. These relatively discrete catalytic sites are produced by suitably applying the transition metal (in an appropriete state) to a substrate, suitably an inorparitie substrate material which can include cathor/graphite.

(D014) The size of the substrate particle is a matter of some importance dependent upon the engineering of the process their. For example, it has fary formation is to take place in a fluid both type of reaction zone, the substrate particle size will suitably be less than about 400 µm. If the fluid both is an ebullient bed of catelyst particles, particle sizes of about 50 to 300 µm have been so found to be preferable. If the fluid bed is an ebullient bed of fortile containing small amounts of catalyst particles, i.e. up to about ten opercent, these should preferably have a size of about 10 to 100 µm. If the fluid bed is a transport bed, either up flow or down flow, the catalyst a carrying particles will confirm the less than about 10 um.

preferably less than about one jum. [0015] It has been found that depositing transition metals according to present claim 1 on small particle substrates produces a catalyst well suitable to use in this 35 invention. The substrate is a material which can conveniently withstand the rigors of Italy flow material which can conveniently withstand the rigors of Italy flow material which can conveniently withstand the rigors of Italy flow material which can conveniently withstand the rigors of Italy flow material which can be substrates include authors, orgaphite, inorqual to the particular transition on the production of t

[0016] Thus, very small particle iron such as might be produced by decomposition of iron compounds, can be deposited on very small particle alumina, e.g., furned alumina having particle sizes on lo larger than about 50 149 jmr (100 mesh). These alumina particles may be made up of individual crystallites which are on the order of about 5 to 20 mm (50 to 200Å), which agglomerate to from particles kning sustaintials available surface a rea sufficient to receive deposits of appropriately sized transition metal catalyst.

[0017] The substrate particles are suitably less than about 300  $\mu m$ . They may be less than 1  $\mu m$  in transport

bed use. It appears that the transition metal reacts with the substrate crystallities such as to bond the metal to the substrate and fix its position, so as to prevent or reterd catalyst agoinmentation, at least for so long as it takes to contact the supported transition metal with the suitable carbon source at appropriate reaction conditions. Upon contact, the carbon source seems to prolyze on the catalytic site and the desirable morphology fibril grows therefrom.

[0018] As noted, the state of the transition metal catalyst stee during froil formation is important to the practice of this invention. Sometimes, it appears that this desirable catalytic site state as well as the state of the substrate carrier therefore is changing during the whole process hereof. Thus, the catalytic sites may applicant et or disperse to some extend during the period from introduction into the reaction zone until the fairlis are ferovered, particles of transition metal catalyst which are sometimes recovered with the fairlis are of 3.5 to 70 nm (35 to 700A), preferably 5 to 30 nm (60 to 300A) in its. Thus, it is believed that the size of the active catalyst steel during fairl formation is substantially comparable to the fairlist are formation.

bit to the diameter of the fibril being formed [0019] It appears that as furtiformation takes place, active catalyst sites become catalytically expended and need to be replaced. Additionally, it has been found that the fibril forming process is more efficient and capable of better comfort in the catalyst is added to the reaction zone intermittently or continuously over substantially the entire course of the reaction, or telest a substantially portion thereof. It is possible that the catalyst containing substate of this invention may ablate with use. That is, when a fibril is formed on a particular catalytic site, that fibril and its associated site may break off from the substrate, with or without some of the substants, thereby exposing further catalytic sites which were previously inside the substrate particular. Thus, periodic or continu-

ous addition of fresh catalyst is desimable.

9 (0020) Thus, according to this invention, the tibril forming process hereof is preferably substantially continuous in that a suitable source of carbon, with or without carrier gas, and catalyst containing particles are continuously or intermittently tell to a reaction zone mainstant at a fibril forming temperature appropriate to the carbon source being used; while Etnip product, usually admixed with the remnants of the catalyst and some-times substants as well, are continuously or intermit-

tently recovered.

90 (0021) The transition metals may be deposited on the substrate by any commonly used technique for accomplishing such deposition. Vapor deposition, sputtering and impregnation may all be suitable. In particular, it has been found to be expeditious to form a water solution or dispersion of the desired metal or metals, mix the water phase with appropriately sized substrate, and then precipitate the metal (3) onto it is substrate, e.g. by

evaporating the water or any other conventional means.

[0022] It is also possibl to deposit the desired transition metals from an organic (as opposed to aqueous) medium. Suitably the transition metals can be dissolved or suppanded in such medium, for example, as an organometallic compound, and then impregnated onto and a into a suitable authoritant or organic carrier medium is removed, feaving behind the impregnated, deposited transition metals.

[0023] After the transition metals are combined with the substate as deviseds, if may be important to treat 10 this combination so as to activate it for this particular catalytic purpose, e.g., by heading it to separate the metals from other ligands, if any, in the deposition compound, it may also be necessary to adjust the size of the prepared catalyst to make it suitable for use in this 15 mineration. Commission of the combination of the property of the pr

[0024] The catalyst for use in this invention may be put on the substants hereof in any form or chemical oxidate or content substants. It may be the exide or have some other ligation state. It may be the exided prior to use, but this is not necessary since the fibril forming needs in is a reducing environment, and thus the transition metal will be reduced during, or immediately prior to, fibril forming as

[0025] Fibrils which are very thin and long, diameters of 3.5 to 70 nm and L/D of up to 100 or more, are produced using these catalysts. These fibrils, as produced by this process, without the necessity of further treat- 30 ment, and without the coproduction of a thermal carbon overcoat, comprise a carbon laver generally concentric about an axis which comprises multiple essentially continuous layers of ordered carbon atoms, which preferebly and usually are crystalline and graphitic. This, as 35 produced, outer layer of ordered carbon atoms often surrounds an inner layer of less ordered carbon atoms. Most preferred products of this invention are high yields of high quality, thin fibrils of appropriate long length having substantially uniform, concentric, substantially continuous, ordered, multiple layers of carbon about an axial (inner core) region, which as a different composition/crystallinity and is preferably hollow. Such fibrils preferably have up to about 100 times, and more greater length than diameter, have diameters of up to 70nm (700 angstroms) and are substantially cylindrical about a substantially hollow core and graphitic as made and without having been treated at higher temperatures than the original fibril manufacturing temperature.

100261 According to one agreed of this invention, operating with catalyst particles as herian est forth, yields of fibrils of greater than about 30 times the weight of transition metal in the catalyst are achievable. In many cases, particularly with mixed transition metals, yields of between 100 and 200 times the weight of transition of metal in the catalyst have been achieved. It has been found that in comparable processes, combinations of transition metal catalysts have sometimes increased

yields by a factor of as much as 2 or even more.

[0027] The following examples illustrate the practice of this invention. By following one or more of these examples, high yields of unique fibrils as above described are produced.

#### Example 1 (comparative)

[0028] A catalyst was prepared using Degussa humed a alumina with an average particle size of about 10mm (100A) and an aggregate mesh size of -100 (149µm), from acetylacetonate was deposited on these alumina particles in a ratio of about 1 part iron, as the acetylacetonate, to 10 parts by weight of alumina. The resultant particle was heated under a hydrogen/ethylene atmosphere under reaction conditions.

[0029] A one (1) inch tube was heated to about 650°C while it was being purged with argon. A mixed flow of hydrogen, at 100 m/tmin, and ethylene, at 200 m/tmin, was feet to the hot tube for the mirutes whereupon cathyl was introduced into the reactor tube. The ethylenenthydrogen mixture was continued through the tubular reactor for 0.5 hours after which the reactor was allowed to cool to room temperature under argon. Hervesting of the fibrilis so produced showed a yield of greater than 30 times the weight of the iron in the cata-

## Example 2

[0030] Into a 3 L. round bottom flask was added 80.08 g of Degussa furned alumina and 285 ml of methanol. The mixture was stirred to produce a thick paste before a solution of 78.26 g (0.194 moles) of ferric nitrate nonahydrate and 4.00 g (0.0123 moles) of molybdenum(VI) oxide bis(2,4-pentanedionate) in 300 ml of methanol (Fe to Mo atom ratio of 94:6) was added slowly. The thick paste which had collected on the sides of the flask was washed down with 65 ml of additional methanol and the mixture was stirred for 1 hour before house vacuum of 1.1 bar (28 in. Hg) was applied while stirring overnight. The purple-tinted solid was placed in a vacuum of 1.1 bar (28 in. Hg) oven at 100°C for 29 hours. A total of 100.7 g of catalyst was obtained. The catalyst was ground and passed through an 80 mesh (177µm) sieve prior to use. Analysis of the catalyst indicated 9.43% by weight iron and 0.99% by weight molybdenum.

(9031) A vertical furnace containing a 2.54cm (1 inch) quart tube with an internal guart wool plug and thermocouple was equilibrated at 650°C under a down flow of 100 ml/min, hydrogen and 200 ml/min, ethylene. Into the tube (onto the quart wool plug) was added 0.1044/hylene. Into g of the above-described catalyst. After 30 min., the hydrogen/ethylene flow was stopped and the oven was allowed to cool to near come temperature. A total of 1.2434 g of fibrilis was harvested for a yield ratio of 126 times the iron weight contain of the catalyst.

#### Example 3

[0032] A sample of catalyst from example 2 (1.6371 p) was placed in a horizontal furnace under argon and was heated to 300°C. After 30 min. at this temperatrure, the 5 furnace was cooled and 1.4460 g of catalyst was recovered (12% wt. loss), having 11.1% by weight iron and 1.2% by weight molybdenum.

[0033] A vertical tube furnace containing a 2.54 cm (1 in.) quartz tube with an internal quartz wool plug and 10 thermocouple was equilibrated at 650°C under a 100 ml/min. down flow of hydrogen and 200 ml/min. down flow of ethylene. Into the hot tube was added 0.1029 g of the catalyst described above. After 30 min., the hydrogen/ethylene flow was stopped and the oven was 15 allowed to cool to near room temperature under argon. A total of 1.3750 g of fibrils was isolated for a weight yield based on theoretical Iron content of 120 times the iran content.

### Example 4

[0034] The vertical tube furnace described in Example 2 was equilibrated at 700°C under the flow of 100 mi/min. hydrogen and 200 mi/min. propane. Onto the 25 Example 9 quartz wool plug was added 0.1041g of catalyst from Example 2. After 30 min. the fuel gases were stopped and the product was cooled under argon. A total of 0.3993 of fibrils was isolated for a weight yield of 41 times the catalyst iron content.

## Example 5

[0035] The procedure of Example 4 was followed at 650°C using 0.1004 g of catalyst from Example 2. A total of 0.3179 g of fibrils was harvested for a weight yield of 34 times the iron content of the catalyst.

## Example 6

[0036] Into a round bottom flask was added 4.25 g of Degussa furned alumina and 30 ml of methanol. The mixture was mechanically stirred while a solution of 4.33 g (10.7 mmol) of ferric nitrate nonahydrate and 0.51 g (1.56 mmol) of molybdenum(VI)oxide bis(2, 4-45 pentanedionate) in 50 ml of methanol was slowly added. The mixture was stirred for 1 hour before the solvent was removed with the aid of a rotary evaporator The resulting damp solid was vacuum dried at 105°C, 1.1 bar (28 in. Hg) for 18 hours. The resulting catalyst was ground and passed through an 177µm (80 mesh) sieve. A total of 5.10 g of catalyst was obtained. Analysis of the catalyst indicated 9.04% by weight iron and 2.18% by weight molybdenum to be present.

[0037] Fibrils were prepared following the procedure 55 of Example 2 at 650°C using 0.0936 g of the above catalyst. A total of 0.9487 g of fibrils was isolated for a weight yield of 126 times the catalyst iron content.

#### Example 7

100381 Into a round bottom flask was added 3.80 g of Degussa furned alumina and 30 ml of methanol. The mixture was mechanically stirred while a solution of 4.33 g (10.7 mmol) of ferric nitrate nonahydrate and 2.04 g (6.25 mmol) of molybdenum(VI)oxide bis(2, 4pentanedionate) in 100 ml of solvent was added. The mixture was held at 105°C and 1.1 ber (28 in. Ho) for 17 hrs. The dried catalyst was sieved (177µm(80 mesh)) to produce 6.10 g of powder. Analysis of the catalyst indicated 8.61% iron and 8.13% molybdenum by weight. [0039] Fibrils were prepared following the procedure of Example 2 at 650°C using 0.1000 g of the above catalyst. A total of 0.8816 g of fibrils was isolated for a weight yield of 102 times the catalyst Iron content.

#### Example 8

20 [0040] The procedure of Example 7 was followed at 700°C using methane and 0,1016g of catalyst. A total of 0.0717g of fibrils was isolated for a yield of 8.2 times the iron content of the catalyst.

[0041] Into a 500 ml round bottom flask was placed 4.37 g of Degussa fumed alumina and 28 ml of methanol. To the stirred mixture was added a solution of 4.33 g (10.7 mmol) of ferric nitrate nonahydrate and 0.46 g (1.32 mmol) of chromium acetylacetonate in 75 ml of methanol. The mixture was stirred for 1 hr. before it was dried for 18 hr. at 105°C and 1.1 bar (28 in. Hg). The catalyst was ground and sieved (177µm(80 mesh)) to produce 5.57 g of powder. The theoretical metal content by weight was 11.9% iron and 1.4% chromium.

[0042] Fibrils were prepared following the procedure of Example 2 at 650°C using 0.0976 g of the above catsiyst. A total of 0.9487 c of fibrils was isolated for a yield 40 of 82 times the theoretical iron content.

## Example 10 (comparative)

(0043) Into a 500 ml round bottom flask was placed 4.40 g of Degussa furned alumina and 35 ml of methanol. To the thick paste was added 4.32g (10.7 mmol) of terric nitrate nonahydrate in 35 ml of methanol. The mixture was stirred for 45 min, before the solid was dried at 95°C and 1.1 bar (28 in. Hg)for 18 hr. The catalyst was ground and sieved(177µm(80 mesh)).

[0044] Fibrils were prepared following the procedure of Example 2 at 650°C using 0.0930 g of the above catalyst. A total of 0.4890 g of fibrils was isolated for a weight yield of 46 times the catalyst iron content.

#### Example 11(comparative)

[0045] Into a round bottom flask was placed 4.33 g of

Degussa furned alumina in 30 mt of methanol. To the strend paste was actided a solution of 4.35 g (1.7 mmol) of ferric noistate nonethydrate and 0.42 g (1.19 mmol) of ferric acetylacetonate in 50 mt of methanol. The mixture was stirred for 75 min. before dryling at 105°C and 1.1 g bar (28 in. Hg) for 17 ms. The solid was ground and served (177µm(0) mesh) to yield 5.87 g of catalyst. Analysis showed 13.79% iron present in the catalyst. [0046] Fibris were prepared following the procedure

[0046] Fibrits were prepared following the procedure of Example 2 at 650°C using 0.0939 g of the above catalyst to produce 0.3962 g of fibrits. This corresponds to 31 times the Iron content of the catalyst.

#### Example 12

(0047) Into a round bottom flask was acided 4.33g of Degussa furned alumina in 20 not water followed by a solution of 4.33 g (10.7 mmol) of ferric nitrate nonety-retate and 0.17g (.138 mmol) of ammonium molybdate in 40 nr of varter. The mixture was mechanically stirred 20 for 1 hour. The water was removed at reduced pressure at 40°C overright. Final drying was accomplished at 140°C and 0.039 bar (26 mm. Hg)for 21 hours to produce 5.57 g of solid. Analysis of the catalyst showed 9.87% by weight iron and 1.45% by weight molybdenum 25 to be present.

[0048] Fibrils were prepared following the procedure of Example 2 at 650°C using 0.0794 g of catalyst to produce 0.8556g of fibrils. This corresponds to 111 times the iron content of the catalyst.

## Example 13

[0049] Into a round bottom flask, containing 4.33 g of Degusas furnod alumina and 30 m1 of methanol, was 30 added a solution of 4.33 g (10.7 mmol) of ferric nitrate nonalhydrate and 0.15 g (0.386 mmol) of ceric intrate in 50 m1 of methanol. An additional 20 m1 of methanol was used to weah all the salls into the flask. The mixture was streed for one hour before the solvent was removed at 4 reduced pressure. The solid was dried at 130°C and 0.400 bar (27 mm Hg)for four days to produce 5.32 grams of catallyst. Analysis of the solid indicated 9.40% iron and 0.89% cerium to be pressent.

[0050] Fibrits were prepared following the procedure of Example 2 at 650°C using 0.0941g of catalyst to produce 0.7552g of fibrits. This corresponds to 88 times the iron content of the catalyst

### Example 14

[0051] Into a round bottom flask was added 4.33g of the Degussa fumed alumins and 20 mt of methanol. Onto the alumina was poured a solution of 4.33g (10.7 mmol) of ferric nitrate and 0.31g (1.22 mmol) of margianese(II) acetylacetonate in 50 ml of methanol. The solvent was removed at reduced pressure of 0.040 bar (27 mm Hg) and the damp solid was vacuum dried at 140°C to promote the control of the description of the control of the damp of the damp solid was vacuum dried at 140°C to pro-

duce 5.18g of solid. Analysis of the catalyst indicated 9.97% iron and 1.18% manganese.

[0052] Fibrils were prepared following the procedure of Example 2 at 650°C using 0.070g of catalyst to produce 0.4948g of fibrils. This corresponds to 66 times the iron content of the catalyst.

#### Example 15

[0053] Into a round bottom flask was added 4,33g of Degussa furned alumina and 30 m of methanol. Onto the atumina was pouved a solution of 4.33g (10.7 mmol) of atumina was pouved a solution of 4.33g (10.7 mmol) of manganese(III) aostylacetonate in 50 m of methanol. The solvent was removed at reduced pressure and the damp solid was reacuum dried at 140°C to produce 5.2°g of solid. Analysis of the catalyst indicated 10.00% iron and 1.18% manganese by weight.

[0054] Fibrils were prepared following the procedure of Example 2 at 650°C using 0.0723g of catalyst to produce 0.7891g of fibrils. This corresponds to 110 times the iron content of the catalyst on a weight basis.

## Example 16 (comparative)

10055] Degussa tumed alumina (400g) and deionized water (8.01) were added to a 22. It flask equipped with a stirrer, pH meter and probe, and two 2 L addition funds. One furned contained an appeaus solution of ferric nitrate nonahydrate (511g dissolved in 5654 ml of water) and the other an appeaus solution of solution to the other an appeaus solution of solution bloatbonate (480g dissolved in 5700 ml of water).

to 6.0 by adding the sodium bloarbonate solution to 5 raise in to the ferrire intrate solution to lower it Next, both solutions were added simultaneously over 3-4 hours with good agliation while maintaining the pH at 6.0. When the addition was complete, stirring was continued for an additional 1/2 hour, after which the slurry was file tered on a 32 cm Buchner funnel. The filter cake was then washed with delonized water and returned to the 22 L flask. Next, additional delonized water was added and the slurry stirred for another 1/2 hour. The batch was then filtered, washed with delonized water, and or succum-drief at 100°C to constant weight (475g). Following drying, the final catalyst was prepared by grinding and sienfront be norduct to 177um (-80 mesh).

## Example 17

[0057] This Example illustrates the practice of this invention using periodic addition of catalyst to produce high fibril yields. A 10.16 cm (four-inch) quartz tube, dosed on the bottom, was placed in a 10.16(4 st inch)diameter x 60.96 cm (24 inch) long furnace. The tube was purged with argon while being heated to 620°C. When the tube was hot, the gas feed was switched to a mixture of hydrogen (1.0 limit) and ethylicities.

ene (5.6 Vmin) via a dip tube to the bottom of the 10.16 cm (4 inch)tube. After 5 min of purging, the catalyst addition was begun.

(0058) A total of 41.13g of catalyst, prepared as described in the Example 16, was added to the hot 5 reactor reservoir. The catalyst was added periodically to the hot reactor in small portions (0.2g) over a period of approximately 6 hours. After catalyst addition was complete, the reaction was allowed to run for an additional one hour and the reactor then cooled to room tempera- 10 ture under argon. The fibrils were removed from the tube and weighed. This batch gave 430g total yield of fibrils which is unusually high for a catalyst based upon iron has the only transition metal. In single batch addition of an iron only catalyst, fibril yields of about 30 15 times the iron content have been observed whereas here the fibril yield is more than 70 times the iron content of the catalyst.

#### Example 18

(0059) The tube and furnace described in Example 17 were heated to 650° under an argon purge. When the tube was not the gas feed was switched to hydrogen and ethylene as described in Example 17.

[0060] A total of 20.4g of catalyst (Fe-Mo) prepared as described in Example 2 was added in a manner similar to that described in Example 17. This batch gave a total fibril vield of 255a.

## Claims

- 1. A continuous process for producing carbon fibrils by decomposing a source of carbon at elevated temperatures in contact with a multivalent metal 35 and recovering the fibrils formed thereby comprising the steps of introducing catalyst particles having a size of up to 400 um and comprising, as the multivalent metal iron mixed with at least one other transition metal on a particulate substrate into a 40 reactor heated to a temperature of 500°C to 1500°C, and recovering the fibrils formed thereby wherein said metal is present on said substrate as a multiplicity of discontinuous catalytic sites which, at least during fibril formation, have a size of 3,5 to 45 70 nm (35 to 700 Å) which size is measured by measuring the size of the transition metal particles recovered along with produced fibrils.
- 2. The process of claim 1 wherein said transistion 50 metal comprises iron with a catalyst site size of about 6 to 30 nm (60 to 300 Å).
- 3. The process of claim 1 wherein said other transition sisting of molybdenum and chromium.
- 4. The process of claim 1 wherein said carbon source

## is a mixture of hydrocarbons

- 5. The process of claim 1 wherein said carbon source is at least one member selected from the group consisting of methane, ethane, propane, butane, benzene, butene, isobutene, cyclohexane, ethylene, propylene, acetylene, toluene, xylene, cumene, ethyl benzene, naphthalene, phenanthrene, anthracene, formaldehyde, acetaldehyde, acetone, methanol, ethanol, and carbon monoxide.
- 6. The process as claimed in claim 1 carried out in an ebulliating bed wherein said catalyst particles have a particle size of about 50 to 300 µm (microns).
- 7. The process of claim 1 carried out in a transport bed wherein said catalyst particles have a particle size of less than about 10 µm (microns).
- 20 8. The process of claim 7 wherein said catalyst particles have a particle size of less than about 1 um (micron).
  - 9. The process of claim 1 including physically cofeeding said catalyst particles and said source of carbon downflow through a reaction zone.
    - 10. The process of claim 1 including physically cofeeding said catalyst particles and said source of carbon upflow through a reaction zone.
    - 11. The process of claim 1 including cafeeding a nonhydrocarbonaceous gas with said carbon contributing feed.
  - 12. The process of claim 1 wherein said catalyst particles are added to the heated reactor over the course of the reaction.
- 13. The process of claim 12 wherein said catalyst particles are added intermittently over the course of the reaction
- 14. The process of claim 1 wherein said source or carbon is added to the heated reactor
- 15. The process of claim 1 wherein said source of carbon is added to the heated reactor over the course of the reaction.
- 16. The process of claim 1 wherein said catalyst particles and said source of carbon are added to the heated reactor over the course of the reaction.
- metal is at least one selected from the group con- 55 17. The process of claim 1 including recovering a high vield of product comprising carbon fibrils characterized as tubes that are substantially free of pyrolytically deposited thermal carbon and have graphitic

layers that are substantially parallel to the fibril axis.

- The process of claim 1 wherein the diameter of said fibrils is between 3.5 and 70 nm. inclusive.
- The process of claim 1 wherein said process is carried out in a fluidized bed.
- 20. A process for continuously producing high yields of long, thin fibrils comprising at least long portions 10 having atomically ordered outer surfaces, as made by contacting a vaporous source of the atoms which will comprise said outer surface with catalyst particles; which process comprises introducing said catalyst particles into a reactor heated to a temperature of 500°C to 1500°C, said catalyst particles having a size of up to 400 um (microns) and comprising, as the multivalent metal iron mixed with at least one other transition metal on a particulate substrate, said metal being present on said sub- 20 strate as a multiplicity of discontinuous catalytic sites which, at least during fibril formation, have a size of 3.5 to 70 nm (35 to 700 Å) which size is measured by measuring the size of the transition metal particles recovered along with produced 25 fibrils, and recovering the fibrils formed thereby.

## Patentansprüche

- 1. Kontinuierliches Verfahren zur Erzeugung von Koh- 30 lenstoff-Fibrillen durch Zersetzen einer Kohlenstoff-Qualle bei erhöhter Temperatur in Gegenwart eines multivalenten Metalls und Wiedergewinnen der dadurch gebildeten Fibrillen, umfassend das Einführen von Katalysator-Teilchen mit einer Größe bis 35 zu 400 µm und umfassend Eisen als multivalentes Metall, das mit zumindest einem anderem Übergangsmetall gemischt ist, auf einem körnigen Substrat in einen Reaktor, der auf eine Temperatur von 500 bis 1500°C erwarmt ist, und Wiedergewinnen 40 der dadurch gebildeten Fibrillen, worin das Metall auf dem Substrat als eine Vielzahl von diskontinuierlichen, katalytischen Stellen vorhanden ist, die zumindest während der Fibrillenbildung eine Größe von 3,5 bis 70 nm (35 bis 700 Å) haben, wobei die 45 Größe durch Messen der Größe der Übergangsmetall-Teilchen gemessen wird, die zusammen mit erzeugten Fibrillen wiedergewonnen werden.
- Verfahren nach Anspruch 1, worin das Übergangsmetall Eisen mit einer Größe der Katalysatorstelle von etwa 6 bis 30 nm (60 bis 300 Å) umfaßt.
- Verfahren nach Anspruch 1, worin das andere Übergangsmetall zumindest eines ist, ausgewählt 55 aus der Gruppe, bestehend aus Molybdän und Chrom

- Verfahren nach Anspruch 1, worin di Kohlenstoff-Quelle eine Mischung aus Kohlenwasserstoffen ist.
- Verlahren nach Anspruch 1, worin die Kohlenstoff-Quelle zumindest eine Verbindung ist, ausgewehlt aus der Gruppe, bestehend aus Methan, Ethan, Propan, Butan, Benzol, Butan, Isobulen, Oyciohezan, Ethylen, Propyten, Acetylen, Toluol, Xylol, Cumol, Ethylbenzol, Naphthalin, Phenarthren, Anthracen, Formatleryd, Acetaldehyd, Aceton, Methanol, Ethanol und Kohlemmonoodi.
- Verfahren nach Anspruch 1, das in einem Siedebett durchgeführt wird, worin die Katalysator-Teilchen eine Teilchengröße von etwa 50 bis 300 μm (Milron) haben.
- Verfahren nach Anspruch 7, worin die Katalysator-Teilchen eine Teilchengröße von weniger als etwa 1 μm (Mikron) haben.
- Verfahren nach Anspruch 1, umfassend das physikalische gemeinsame Führen der Katalysator-Teilchen und der Kohlenstoff-Quelle im Abwärtsstrom durch eine Reaktionszone.
- Verfahren nach Anspruch 1, umfassend das gemeinsame physikalische Führen der Katalysator-Teilchen und der Kohlenstoff-Queille im Aufwärtsstrom durch eine Reaktionszone.
- Verfahren nach Anspruch 1, umfassend das gemeinsame Führen eines nicht-Kohlenwasserstoft-haltigen Gases mit der Kohlenstoff-haltigen Zufuhr.
- Verfahren nach Anspruch 1, worin die Katalysator-Teilchen zu dem erwärmten Reaktor w\u00e4hrend der Reaktion zugegeben werden.
- Verfahren nach Anspruch 12, worin die Katalysator-Teilchen absatzweise w\u00e4hrend des Verlaufes der Reaktion zugegeben werden.
- 50 14. Verfahren nach Anspruch 1, worin die Kohlenstoff-Quelle zu dem Heizreaktor gegeben wird.
  - Verfahren nach Anspruch 1, worin die Kohlenstoff-Quelle zu dem erwärmten Reaktor während des Verlaufes der Reaktion gegeben wird.
  - Verfahren nach Anspruch 1, worin die Katalysator-Teilchen und die Kohlenstoff-Quelle zu dem Heizre-

aktor während des Verlaufes der Reaktion gegeben werden.

- 17. Verfahren nach Anspruch 1, umfassend das Wiedergewinnen einer hohen Ausbeute des Produktes, 5 umfassend Kohlenstoff-Fibrillen, gekennzeichnet als Rohre, die im wesentlichen frei sind von pyrobysisch niedergeschlagenen, thermischern Kohlenstoff und graphtische Schichten haben, die im wesentlichen narallel zu der Fibrillense sind.
- Verfahren nach Anspruch 1, worin der Durchmesser der Fibrillen zwischen 3,5 und 70 nm einschließlich ist
- Verfahren nach Anspruch 1, worin das Verfahren in einem Fließbett durchgeführt wird.
- 20. Verfahren zum kontinuierlichen Erzeugen hoher Ausbeuten von langen, dünnen Fibrillen, umfas- 20 send zumindest lange Bereiche mit atomisch angeordneten äußeren Oberflächen, erzeugt durch Kontaktieren einer dampfartigen Quelle der Atome. die die außere Oberfläche umfassen, mit Katalysator-Teilchen; wobei das Verfahren das Einführen 25 von Katalysator-Teilchen in einen Reaktor, der auf eine Temperatur von 500 bis 1500°C erwarmt ist, wobei die Katalysator-Teilchen eine Größe von bis zu 400 um (Mikron) haben und als multivalentes Metall Eisen, das mit zumindest einem anderen 30 Übergangsmetall vermischt ist, auf einen körnigen Substrat enthalten, wobei das Metall auf dem Substrat als eine Vielzahl von diskontinuierlichen katalytischen Stellen vorhanden ist, die zumindest während der Fibrill-Bildung eine Größe von 3,5 bis 35 70 nm (35 bis 700 Å) haben, wobei die Größe durch Messen der Größe der Übergangsmetall-Teilchen gemessen wird, die zusammen mit den erzeugten Fibrillen wiedergewonnen werden, und das Wiedergewinnen der dadurch gebildeten Fibrillen umfaßt.

## Revendications

1. Procédé en continu destiné à la production de tibrilles de carbone par décomposition d'une source et carbone à des températures élevées en contact avec un métal plurvalent et lécupération des fibrilles unies formées, comprenant les étapes consistant à introduire des particules de catalyseur présentant une dimension alant jusqu'à 400 µm et so comprenant, en tant que métal plurvalent, du fer métal de sa un mois un autre métal de s'amsion sur un substrat particulaire dans un réscriter charifé à un moins un autre métal de s'amsion sur un substrat particulaire dans un réscriter charifé à un empérature de 500° C à 150° C, et à récupérer les fibrilles ains formées, dans lequel s'electif métal est présent sur lett substrat sous forme d'une multiplicité de sites catalyfiques discontinus lesquels, au moins pendant la formation des

fibrilles, présentent une dimension de 3,5 à 70 nm (35 à 700 Å), laquelle dimension est mesurée en mesurant la dimension des perticules de métal de transition récupérées en même temps que les fibrilles produites.

- Procédé selon la revendication 1, dans lequel ledit métal de transition comprend du ser présentant une dimension de sites de catalyseur d'environ 6 à 30 nm (60 à 300 Å).
- Procédé selon la revendication 1, dans lequel ledit autre métal de transition est au moins l'un choisi parmi le groupe constitué du molybdène et du chrome.
- Procédé selon la revendication 1, dans lequel ladite source de carbone est un mélange d'hydrocarbures.
- 5. Procedé selon la revendication 1, dans lequel lastite source de carbone est un mois un elément choisi parmi le groupe constituté du méthane, de l'éthane, du propane, du butane, du butane, du butane, du la l'alcohutène, du cyclohexane, de l'éthytène, du propane, de l'éthytène, du propiène, de l'activiténe, du louiene, du syène, du cumène, de l'éthytènenaire, du naphtalène, du prinamitréne, de l'arthréchen, du formétisfyide, de l'actéstidéne, de l'arthréchen, du formétisfyide, de l'actéstidés de carbone.
  - Procédé selon la revendication 1, exécuté dans un it à ébuilition dans lequel lesdites particules de catalyseur présentent une dimension de particules d'environ 50 à 300 µm (microns).
- Procédé selon la revendication 1, exécuté dans un lit à transport dans lequel lescrites particules de catalyseur présentent une dimension de particules de moins d'environ 10 um (microns).
  - Procédé selon la revendication 7, dans lequel lesdites particules de catalyseur présentent une dimension de particules de moins d'environ 1 μm (microns).
  - Procédé selon la revendication 1, comprenant une coalimentation physique desdites particules de catalyseur et de ladite source de carbone en circulation vers le bas à travers une zone réactionnelle.
- 10. Procédé selon la revendication 1, comprenant une coalimentation physique desdites particules de catalyseur et de ladite source de carbone en circulation vers le haut à travers une zone réactionnelle.
- Procédé selon la revendication 1, comprenant une coalimentation d'un gaz non-hydrocarboné avec

ladite alimentation de contribution en carbone.

- Procédé selon la revendication 1, dans lequel lesdites particules de catalyseur sont ajoutées au réacteur chauffé pendant la durée de la réaction.
- Procédé selon la revendication 12, dans lequel lesdites particules de catalyseur sont ajoutées par intermittence pendant la durée de la réaction.
- Procédé selon la revendication 1, dans lequel ladite source de carbone est ajoutée au réacteur chauffé.
- Procédé selon la revendication 1, dans lequel ladite source de carbone est ajoutée au réacteur chauffé 15 pendant le durée de la réaction.
- Procédé selon la revendication 1, dans lequel lesdites particules de catalyseur et ladite source de carbone sont gioutées au réacteur chauffé pendant la 20 durée de la réaction.
- 17. Procédé selon la revendication 1, comprenant la récupération d'un rendement élevé de produit comprenant des fibrilles de carbone caractérisées 32 comme étant des tabre qui sont prasquement exempts de carbone thermique déposé de tapon pyrolyfolue et comportent des couches graphiliques qui sont pratiquement paraltèles à l'axe des fibrilles.
- Procédé selon la revendication 1, dans lequel le diamètre desdites fibrilles est entre 3,5 et 70 nm, inclus.
- Procédé selon la revendication 1, dans lequel ledit procédé est exécuté dans un lit fluidisé.
- 20. Procédé destiné à produire en continu des rendements élevés de fibrilles longues, fines, compre- 40 nant au moins des parties longues présentant des surfaces externes atomiquement ordonnées, réalisées en mettant en contact une source sous forme de vapeur des atomes qui constitueront ladite surface externe avec des particules de catalyseur, lequel procédé comprend l'introduction desdites particules de catalyseur dans un réacteur chauffé à une température de 500° C à 1 500° C, lesdites particules de catalyseur présentant une dimension allant jusqu'à 400 µm (microns) et comprenant, en 50 tant que métal plurivalent, du fer mélangé à su moins un autre métal de transition sur un substrat particulaire, ledit métal étant présent sur ledit substrat sous forme d'une multiplicité de sites catalytiques discontinus lesquels, au moins pendant la 55 formation des fibrilles, présentent une dimension de 3,5 à 70 nm (35 à 700 Å), laquelle dimension est mesurée en mesurant la dimension des particules

de métal d transition récupérées en même temps que les fibrilles produites, et la récupération des fibrilles ainsi formées.

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